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APPLICATION NO.	FILING DATE	FIRST NAMED INVENTOR	ATTORNEY DOCKET NO.	CONFIRMATION NO.
10/618,125	07/11/2003	James D.B. Smith	2003P10020US	4105
7590 . 06/22/2005		EXAMINER		
Siemens Corporation			FEELY, MICHAEL J	
Intellectual Prop	erty Department		· · · · · · · · · · · · · · · · · · ·	
170 Wood Avenue South			ART UNIT	PAPER NUMBER
Iselin, NJ 08830			1712	
	·		DATE MAILED: 06/22/2005	

Please find below and/or attached an Office communication concerning this application or proceeding.

	Application No.	Applicant(s)				
Office Action Summers	10/618,125	SMITH, JAMES D.B.				
Office Action Summary	Examiner	Art Unit				
7	Michael J. Feely	1712				
The MAILING DATE of this communication ap Period for Reply	pears on the cover sheet with t	the correspondence address				
A SHORTENED STATUTORY PERIOD FOR REPL THE MAILING DATE OF THIS COMMUNICATION. - Extensions of time may be available under the provisions of 37 CFR 1. after SIX (6) MONTHS from the mailing date of this communication. - If the period for reply specified above is less than thirty (30) days, a replet if NO period for reply is specified above, the maximum statutory period Failure to reply within the set or extended period for reply will, by statut Any reply received by the Office later than three months after the mailing earned patent term adjustment. See 37 CFR 1.704(b).	136(a). In no event, however, may a reply oly within the statutory minimum of thirty (3 will apply and will expire SIX (6) MONTHS te, cause the application to become ABANI	be timely filed D) days will be considered timely. From the mailing date of this communication. DONED (35 U.S.C. § 133).				
Status						
1) Responsive to communication(s) filed on 15 A	April 2005.					
	s action is non-final.					
	Since this application is in condition for allowance except for formal matters, prosecution as to the merits is closed in accordance with the practice under <i>Ex parte Quayle</i> , 1935 C.D. 11, 453 O.G. 213.					
Disposition of Claims	Lx parte Quayle, 1935 C.D. 1	1, 400 O.G. 215.				
<u> </u>						
 4) Claim(s) 1-22 is/are pending in the application. 4a) Of the above claim(s) is/are withdrawn from consideration. 5) Claim(s) is/are allowed. 6) Claim(s) 1,2,4-10 and 12-22 is/are rejected. 7) Claim(s) 3 and 11 is/are objected to. 8) Claim(s) are subject to restriction and/or election requirement. 						
Application Papers						
 9) The specification is objected to by the Examination 10) The drawing(s) filed on 11 July 2003 is/are: a Applicant may not request that any objection to the Replacement drawing sheet(s) including the correction 11) The oath or declaration is objected to by the Examination) accepted or b) objected or b) objected or b) objected or abeyance. otion is required if the drawing(s)	See 37 CFR 1.85(a). s objected to. See 37 CFR 1.121(d).				
Priority under 35 U.S.C. § 119						
12) Acknowledgment is made of a claim for foreign a) All b) Some * c) None of: 1. Certified copies of the priority documen 2. Certified copies of the priority documen 3. Copies of the certified copies of the priority documen application from the International Burea * See the attached detailed Office action for a list	nts have been received. Its have been received in Applority documents have been received in the control of the	ication No ceived in this National Stage				
Attachment(s)						
1) Notice of References Cited (PTO-892) 4) Interview Summary (PTO-413)						
2) Notice of Draftsperson's Patent Drawing Review (PTO-948) 3) Information Disclosure Statement(s) (PTO-1449 or PTO/SB/08 Paper No(s)/Mail Date	Paper No(s)/M	ail Date mal Patent Application (PTO-152)				

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DETAILED ACTION

Pending Claims

Claims 1-22 are pending.

Claim Objections

1. The objection to claims 1-13 has been overcome by amendment.

Claim Rejections - 35 USC § 103

- 2. The following is a quotation of 35 U.S.C. 103(a) which forms the basis for all obviousness rejections set forth in this Office action:
 - (a) A patent may not be obtained though the invention is not identically disclosed or described as set forth in section 102 of this title, if the differences between the subject matter sought to be patented and the prior art are such that the subject matter as a whole would have been obvious at the time the invention was made to a person having ordinary skill in the art to which said subject matter pertains. Patentability shall not be negatived by the manner in which the invention was made.
- 3. The rejection of claims 1, 2, 4-10, 12, 13, and 18-22 under 35 U.S.C. 103(a) as being unpatentable over Campbell et al. (Pub. No. 2004/0102529) stands for the reasons of record.

Regarding claims 1, 2, 4, 6-8, 10, and 13, Campbell et al. disclose: (1) a method of making homogeneous LCT-epoxy polymers with HTC-oligomers (Abstract; paragraphs 0012 and 0016) comprising: grafting at least one functionalized organic group onto at least one nanosized HTC-material to produce HTC oligomer product (Abstract; paragraphs 0025-0028); reacting said HTC-oligomer product with at least on LCT-epoxy resin under sufficient conditions to form a uniform dispersion and an essentially complete co-reactivity of said HTC-oligomer product with said at least one LCT-epoxy resin, wherein a mixture is formed (paragraphs 0029-0033); and curing said mixture to produce said homogeneous LCT-epoxy polymers with HTC-oligomers (paragraphs 0029-0033);

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- (2) wherein said at least one nano-sized HTC-material comprises at least one of alumina, silica and a metal oxide (Abstract; paragraph 0025);
- (4) wherein the grafting of said at least one organic group onto said at least one nano-sized HTC-material is performed by at least one of a silane grafting and a free radical grafting (Abstract; paragraphs 0025-0028);
- (6) wherein reacting said HTC-oligomer product with said at least one LCT-epoxy further comprises warming until said mixture is clear (paragraphs 0028-0029);
- (7) further comprising mixing at least one anhydriding agent with at least one of said at least one LCT-epoxy resin and said HTC-oligomer product, wherein said homogeneous LCT-epoxy polymers with HTC-oligomers are a homogeneous LCT-epoxy anhydride polymers with HTC-oligomers (paragraphs 0033-0034); (8) wherein said anhydriding agent is taken from the group consisting of 1-methylhexahydrophthalic anhydride and 1-methyltetrahydrophthalic anhydride (paragraph 0034);
- (10) further comprising mixing at least one vinyl agent with at least one of said at least one LCT-epoxy resin and said HTC-oligomer product, wherein said homogeneous LCT-epoxy polymers with HTC-oligomers are a homogeneous LCT-epoxy vinyl polymers with HTC-oligomers (paragraph 0043); and
- (13) wherein said mixture is added to an electrical insulator as a coating before curing (paragraphs 0048-0049).

Campbell et al. are deficient in that they do not explicitly disclose: (1) a ratio of HTC-oligomer to LCT-epoxy resin of between 1:4 and 3:1 (20-75%) by weight; and (2) an overall dielectric strength of at least 1.2 kV/mil.

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Regarding (1) the ratio of HTC-oligomer to LCT-epoxy resin, Campbell et al. add colloidal silica, functionalized with organoalkoxysilane, to curable epoxy formulations (including biphenyl type epoxy system) in order to provide a formulation with a low viscosity prior to curing and a low coefficient of thermal expansion after curing (see paragraph 0012). The functionalized silica acts in place of traditional fillers in order to enhance the physical properties of epoxy formulation. As with all filler/resin systems, the quantity of the filler is adjusted in order to optimize flow/structural properties, along with overall physical and/or electrical properties of the cured product. Hence, the proportion of HTC-oligomer and LCT-epoxy resin is a result effective variable. It should also be noted that applicant fails to demonstrate criticality for this claimed range.

In light of this, it has been found that, "[W]here the general conditions of a claim are disclosed in the prior art, it is not inventive to discover the optimum or workable ranges by routine experimentation." – *In re Aller*, 220 F.2d 454, 456, 105 USPQ 233, 235 (CCPA 1955); and, "A particular parameter must first be recognized as a result-effective variable, i.e., a variable which achieves a recognized result, before the determination of the optimum or workable ranges of said variable might be characterized as routine experimentation." – *In re Boesch*, 617 F.2d 272, 205 USPQ 215 (CCPA 1980).

Therefore, it would have been obvious to one of ordinary skill in the art at the time of the invention to use the claimed ratio range of 1:4 to 3:1 of HTC-oligomer to LCT-epoxy in the composition of Campbell et al. because it has been found that where the general conditions of a result effective variable are disclosed in the prior art, it is not inventive to discover the optimum or workable ranges by routine experimentation.

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Regarding (2) an overall dielectric strength of at least 1.2 kV/mil, it appears that this property would have been an inherent feature in Campbell et al. because this obvious composition satisfies the chemical requirements of the instant invention. It has been found that, "Products of identical chemical composition can not have mutually exclusive properties." A chemical composition and its properties are inseparable. Therefore, if the prior art teaches the identical chemical structure, the properties applicant discloses and/or claims are necessarily present. *In re Spada*, 911 F.2d 705, 709, 15 USPO2d 1655, 1658 (Fed. Cir. 1990).

Regarding claim 5, the teachings of Campbell et al. are as set forth above and incorporated herein. As discussed above, the proportion of HTC-oligomer and LCT-epoxy resin is a result effective variable, and applicant fails to demonstrate criticality for this claimed range.

Therefore, it would have been obvious to one of ordinary skill in the art at the time of the invention to use the claimed proportion of 20-50 wt% of HTC oligomers in the composition of Campbell et al. because it has been found that where the general conditions of a claimed result effective variable are disclosed in the prior art, it is not inventive to discover the optimum or workable ranges by routine experimentation.

Regarding claim 9, Campbell et al. do not explicitly disclose a concentration of 20-40 wt% of anhydriding agent; however, applicant fails to show criticality for this range.

Furthermore, one skilled in the art would have readily recognized that the quantity of hardener in an epoxy resin composition is a result effect variable that ensures adequate curing of the resin system.

Therefore, it would have been obvious to one of ordinary skill in the art at the time of the invention to use the claimed range of 20-40 wt% of anhydriding agent in the composition of

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Campbell et al. because it has been found that where the general conditions of a claimed result effective variable are disclosed in the prior art, it is not inventive to discover the optimum or workable ranges by routine experimentation.

Regarding claim 12, Campbell et al. do not explicitly disclose a concentration of 4-16 wt% of vinyl agent; however, applicant fails to show criticality for this range. Furthermore, it should be noted that Campbell et al. use this vinyl agent as a reactive diluent to decrease viscosity of the overall formulation. Hence, the quantity of this reactive diluent is a result effective variable.

Therefore, it would have been obvious to one of ordinary skill in the art at the time of the invention to use the claimed range of 4-16 wt% of anhydriding agent in the composition of Campbell et al. because it has been found that where the general conditions of a claimed result effective variable are disclosed in the prior art, it is not inventive to discover the optimum or workable ranges by routine experimentation.

Regarding claims 18-22, the teachings of Campbell et al. are as set forth above and incorporated herein. The only limitations not discussed above are:

- (1) a thermal conductivity in the transverse direction of at least 0.50 W/mK and in the thickness direction of at least 0.99 W/mK in an environment of 25°C; and
- (2) wherein said homogeneous LCT-epoxy polymers with HTC-oligomers are substantially free of particle wetting and micro-void formation.

It appears that these properties would have been inherent features in Campbell et al.
because this obvious composition satisfies the chemical requirements of the instant invention. It
has been found that, "Products of identical chemical composition can not have mutually

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exclusive properties." A chemical composition and its properties are inseparable. Therefore, if the prior art teaches the identical chemical structure, the properties applicant discloses and/or claims are necessarily present. *In re Spada*, 911 F.2d 705, 709, 15 USPQ2d 1655, 1658 (Fed. Cir. 1990)

4. Claims 14-17 are rejected under 35 U.S.C. 103(a) as being unpatentable over Campbell et al. (Pub. No. 2004/0102529) in view of Smith et al. (US Pat. No. 6,384,152).

Regarding claims 14-17, the teachings of Campbell et al. are as set forth above and incorporated herein. The only limitations not discussed above are: (14) impregnating said mixture onto an electrical insulator; (17) wherein said electrical insulator is a mica/glass insulating tape.

As mentioned above, the advantage of the composition taught by Campbell et al. is that it provides an epoxy formulation with a low viscosity before cure and a cured product that has a low coefficient of thermal expansion (see paragraph 0012). Specifically, the desired viscosity is between about 50-100,000 cps @ 25°C (see paragraph 0012). In addition, Campbell et al. disclose that their composition is useful in numerous electrical applications (see paragraphs 0048-0049); however, they do not explicitly disclose an impregnated mica/glass insulating tape.

Smith et al. disclose an epoxy resin composition useful for impregnating mica/glass insulating tapes. They disclose that there is a need for an insulating epoxy resin system with a low viscosity that makes it suitable for vacuum impregnation (see column 1, lines 50-63).

Specifically, a viscosity of 10-150 cps @ 25°C is desired (see column 2, lines 9-31).

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In light of this, the resin system of Campbell et al. appears to be suitable for mica/tape impregnations because its viscosity range overlaps the desired viscosity range set forth in Smith et al.

Therefore, it would have been obvious to one of ordinary skill in the art at the time of the invention to incorporate the resin system of Campbell et al. in the mica/glass tape of Smith et al. because the resin system of Campbell et al. provides an epoxy formulation with a low viscosity before cure and a cured product that has a low coefficient of thermal expansion.

Allowable Subject Matter

- 5. Claims 3 and 11 are objected to as being dependent upon a rejected base claim, but would be allowable if rewritten in independent form including all of the limitations of the base claim and any intervening claims.
- 6. The following is a statement of reasons for the indication of allowable subject matter:

 *Regarding claim 3, Campbell et al. is the closest prior art; however, they provide no motivation to substitute their silica for magnesium oxide.

Regarding claim 11, Campbell et al. is the closest prior art; however, they provide no motivation to use p-vinylphenylglycidylether as their vinyl agent.

Response to Arguments

7. Applicant's arguments filed April 15, 2005 have been fully considered but they are not persuasive.

Argument # 1: Campbell, in fact states that the preferred epoxy resins are aliphatic and cycloaliphatic, which would produce a resin with a poor thermal stability, and therefore teaches away from the present invention.

Applicant is reminded that a reference is considered as a whole – both preferred and non-preferred embodiments constitute the prior art. Campbell et al. do prefer aliphatic and cycloaliphatic epoxy resins; however, they also teach the use of aromatic epoxy resin (see paragraph 0016). Included are resins featuring mesogenic structures, which would have qualified as LCT resins.

Argument # 2: In fact, Campbell does not disclose grafting at all. As described in paragraph 26 of Campbell, using organoalkoxysilane is a condensation reaction not a grafting reaction of the present invention. Although applicant teaches silane grafting, one of ordinary skill in the art would know that this is not the same thing as the organoalkoxysilane reaction disclosed in Campbell.

Applicant fails to disclose why their "grafting" is different from the functionalization step set forth in Campbell et al. The "grafting" procedure set forth in the instant Specification (see paragraphs 048-050) is the same as the functionalization step in Campbell et al., wherein hydrolysable groups of the organoalkoxysilanes are hydrolyzed. These hydrolyzed groups are attached silica particles (HTC-material) via a condensation reaction, binding the oxygen atom of the organoalkoxysilane to the surface of the silica particle (HTC-material). By doing this, the

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organofunctional group remains free to create a functionalized silica material. Whether you call it "grafting" or "condensing", it is the same thing. The Examiner would welcome an explanation as to why the claimed "grafting" process is different; however, "one of ordinary skill in the art would know that this is not the same thing as the organoalkoxysilane reaction in Campbell," is not persuasive.

Argument # 3: The examples given in Campbell are on the atomic scale. For instance, the formula shown after paragraph 26 is one atom of silicon in a molecule. Applicant's representation of the HTC material is a nano-sized particle, which comprises many atoms.

The formula after paragraph 26 is the chemical representation of the organoalkoxysilane used to functionalize the colloidal silica. Applicant is directed to paragraph 25, which clearly states that the colloidal silica to be functionalized is a nano-sized particle.

Argument # 4: Applicant would also like to point out that the teachings of Campbell would not produce a polymer with a dielectric strength of at least 1.2 kV/mil. In addition, Applicant is unable to find in Campbell any teachings of what percentage of the polymer is an HTC material.

Applicant fails to say why the teachings of Campbell et al. would not produce a polymer with a dielectric strength of at least 1.2 kV/mil. Furthermore, both of these issues are addressed in the above rejections – see discussion of claims 1, 2, 4, 6-8, 10, and 13.

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Conclusion

8. THIS ACTION IS MADE FINAL. Applicant is reminded of the extension of time policy as set forth in 37 CFR 1.136(a).

A shortened statutory period for reply to this final action is set to expire THREE MONTHS from the mailing date of this action. In the event a first reply is filed within TWO MONTHS of the mailing date of this final action and the advisory action is not mailed until after the end of the THREE-MONTH shortened statutory period, then the shortened statutory period will expire on the date the advisory action is mailed, and any extension fee pursuant to 37 CFR 1.136(a) will be calculated from the mailing date of the advisory action. In no event, however, will the statutory period for reply expire later than SIX MONTHS from the mailing date of this final action.

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Communication

Any inquiry concerning this communication or earlier communications from the examiner should be directed to Michael J. Feely whose telephone number is 571-272-1086. The examiner can normally be reached on M-F 8:30 to 5:00.

If attempts to reach the examiner by telephone are unsuccessful, the examiner's supervisor, Randy Gulakowski can be reached on 571-272-1302. The fax phone number for the organization where this application or proceeding is assigned is 703-872-9306.

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Michael J. Feely Primary Examiner Art Unit 1712